

<https://helda.helsinki.fi>

---

## A coherent feed-forward loop drives vascular regeneration in damaged aerial organs of plants growing in a normal developmental context

Radhakrishnan, Dhanya

2020-03

---

Radhakrishnan , D , Shanmukhan , A P , Kareem , A , Aiyaz , M , Varapparambathu , V , Toms , A , Kerstens , M , Valsakumar , D , Landge , A N , Shaji , A , Mathew , M K , Sawchuk , M G , Scarpella , E , Krizek , B A , Efroni , I , Mähönen , A P , Willemsen , V , Scheres , B & Prasad , K 2020 , ' A coherent feed-forward loop drives vascular regeneration in damaged aerial organs of plants growing in a normal developmental context ' , Development , vol. 147 , no. 6 , 185710 . <https://doi.org/10.1242/dev.185710>

---

<http://hdl.handle.net/10138/317516>

<https://doi.org/10.1242/dev.185710>

---

cc\_by

publishedVersion

---

*Downloaded from Helda, University of Helsinki institutional repository.*

*This is an electronic reprint of the original article.*

*This reprint may differ from the original in pagination and typographic detail.*

*Please cite the original version.*

# A coherent feed-forward loop drives vascular regeneration in damaged aerial organs of plants growing in a normal developmental context

Dhanya Radhakrishnan<sup>1,\*</sup>, Anju Pallipurath Shanmukhan<sup>1,\*</sup>, Abdul Kareem<sup>1,\*</sup>, Mohammed Aiyaz<sup>1,\*</sup>, Vijina Varapparambathu<sup>1</sup>, Ashna Toms<sup>1</sup>, Merijn Kerstens<sup>2</sup>, Devisree Valsakumar<sup>1</sup>, Amit N. Landge<sup>1</sup>, Anil Shaji<sup>1</sup>, Mathew K. Mathew<sup>1,3</sup>, Megan G. Sawchuk<sup>4</sup>, Enrico Scarpella<sup>4</sup>, Beth A. Krizek<sup>5</sup>, Idan Efroni<sup>6</sup>, Ari Pekka Mähönen<sup>7,8</sup>, Viola Willemsen<sup>2</sup>, Ben Scheres<sup>2</sup> and Kalika Prasad<sup>1,†</sup>

## ABSTRACT

Aerial organs of plants, being highly prone to local injuries, require tissue restoration to ensure their survival. However, knowledge of the underlying mechanism is sparse. In this study, we mimicked natural injuries in growing leaves and stems to study the reunion between mechanically disconnected tissues. We show that *PLETHORA* (*PLT*) and *AINTEGUMENTA* (*ANT*) genes, which encode stem cell-promoting factors, are activated and contribute to vascular regeneration in response to these injuries. *PLT* proteins bind to and activate the *CUC2* promoter. *PLT* proteins and *CUC2* regulate the transcription of the local auxin biosynthesis gene *YUC4* in a coherent feed-forward loop, and this process is necessary to drive vascular regeneration. In the absence of this *PLT*-mediated regeneration response, leaf ground tissue cells can neither acquire the early vascular identity marker *ATHB8*, nor properly polarise auxin transporters to specify new venation paths. The *PLT*-*CUC2* module is required for vascular regeneration, but is dispensable for midvein formation in leaves. We reveal the mechanisms of vascular regeneration in plants and distinguish between the wound-repair ability of the tissue and its formation during normal development.

**KEY WORDS:** Vascular regeneration, *PLT*, *CUC2*, Wound repair, Auxin, *PIN1*, *Arabidopsis*

## INTRODUCTION

Plants are prone to numerous injuries in their lifespan, owing to their sessile lifestyle. They are subjected to injuries caused by biotic factors such as pathogen attack and herbivory. Abiotic factors such as damaging weather conditions can also cause tissue damage.

Unhealed wounds can compromise plant fitness and survival, and tissue-healing mechanisms have evolved to counteract the damage. Following wounding, regenerative responses may be restricted to local healing in the form of cell proliferation or may entail complete regeneration of damaged tissue or organ (Ikeuchi et al., 2016; Galliot et al., 2017). The capacity of plants to regenerate the complete body plan *in vitro* from excised tissue is a powerful demonstration of the versatility of plant regeneration processes and forms the basis for many horticultural applications (Kareem et al., 2015; Ikeuchi et al., 2016; Radhakrishnan et al., 2018).

In stem, cellular, molecular and hormonal interactions at wound sites coordinate wound healing and restore vasculature (Flaishman et al., 2003; Asahina et al., 2011; Pitaksaringkam et al., 2014; Melnyk et al., 2015; Mazur et al., 2016). Auxin is important for vascular tissue regeneration in multiple plant species (Sachs, 1968, 1969, 1981, 1991). The canalization models that underlie this regeneration process rely on the potential of auxin to induce correctly polarised auxin transporters together with activation of vascular cell fate determinants (Wenzel et al., 2007; Donner et al., 2009; Ohashi-Ito et al., 2013). In the growing tips of shoots and roots, damaged meristematic cells are replaced using positional cues from neighbouring cells (van den Berg et al., 1995; Reinhardt et al., 2003).

In roots, regeneration involves reactivation of embryo-specific genes, proper reallocation of root cell-fate determinants and integration of auxin, cytokinin and jasmonate signals (Xu et al., 2006; Efroni et al., 2016; Marhava et al., 2019; Zhou et al., 2019).

Laser ablation and root tip resection studies have shown that stem cell activation is a vital step for regeneration of lost cells and entire organs (van den Berg et al., 1995; Xu et al., 2006; Marhava et al., 2019; Zhou et al., 2019). The stem cell regulators *PLETHORA1* (*PLT1*) and *PLT2* are essential for the re-establishment of quiescent centre (QC) cells upon laser ablation and for the regeneration of primary and lateral root tips following resection (Xu et al., 2006; Durgaprasad et al., 2019). *PLT1* and *PLT2* are induced by *PLT3*, *PLT5* and *PLT7* activity to regulate stem cell activation during lateral root development (Du and Scheres, 2017). In the shoot, members of the *PLT* family along with the transcription factor *AINTEGUMENTA* (*ANT*) regulate the development and phyllotaxis of aerial organs (Prasad et al., 2011; Krizek, 2015). *PLT* factors also regulate hormone-mediated *de novo* shoot regeneration (Kareem et al., 2015).

Although several studies have addressed specific regeneration processes in specific plant parts or in excised organs and have implicated certain factors regulating these processes, our knowledge of the underlying molecular mechanisms of wound repair in aerial organs is limited (Ikeuchi et al., 2018). It is largely unknown how

<sup>1</sup>School of Biology, Indian Institute of Science Education and Research, Thiruvananthapuram 695551, India. <sup>2</sup>Plant Developmental Biology, Wageningen University Research, Wageningen 6708 PB, The Netherlands. <sup>3</sup>National Centre for Biological Sciences, Tata Institute of Fundamental Research, 15, Bengaluru, 560065, India. <sup>4</sup>Department of Biological Sciences, University of Alberta, Edmonton, Alberta T6G 2E9, Canada. <sup>5</sup>Department of Biological Sciences, University of South Carolina, Columbia, SC 29208, USA. <sup>6</sup>The Robert H. Smith Institute of Plant Sciences and Genetics in Agriculture, The Hebrew University, Rehovot 76100, Israel. <sup>7</sup>Institute of Biotechnology HiLIFE, University of Helsinki, 00014 Helsinki, Finland. <sup>8</sup>Organismal and Evolutionary Biology Research Programme, Faculty of Biological and Environmental Sciences, Viikki Plant Science Centre, University of Helsinki, 00014 Helsinki, Finland.

\*These authors contributed equally to this work

†Author for correspondence (kalika@iisertvm.ac.in)

DOI: B.A.K., 0000-0001-5821-0180; I.E., 0000-0002-0219-8685; K.P., 0000-0001-8666-856X

wound repair in leaf tissue relates to the normal developmental programme. Here, we investigate vascular reprogramming after leaf damage from the viewpoint that tissue reprogramming may require stem cell factors identified in other regeneration contexts. We reveal an essential role of members of the *PLETHORA* (*PLT*)/*AINTEGUMENTA* (*ANT*) gene family in activating regeneration responses. *PLT* genes act through *CUP-SHAPED COTYLEDON2* (*CUC2*) to repair wounds and regenerate vascular tissue in damaged aerial organs. Furthermore, we show that the *PLT-CUC2* module acts through local auxin biosynthesis, and is required for proper repolarisation of PIN1 auxin efflux facilitators and reprogramming of vascular identity in aerial organs. The *PLT-CUC2* module is strictly required for regeneration of leaf vasculature, but is not essential for the normal development of closed vein loops in the absence of perturbations.

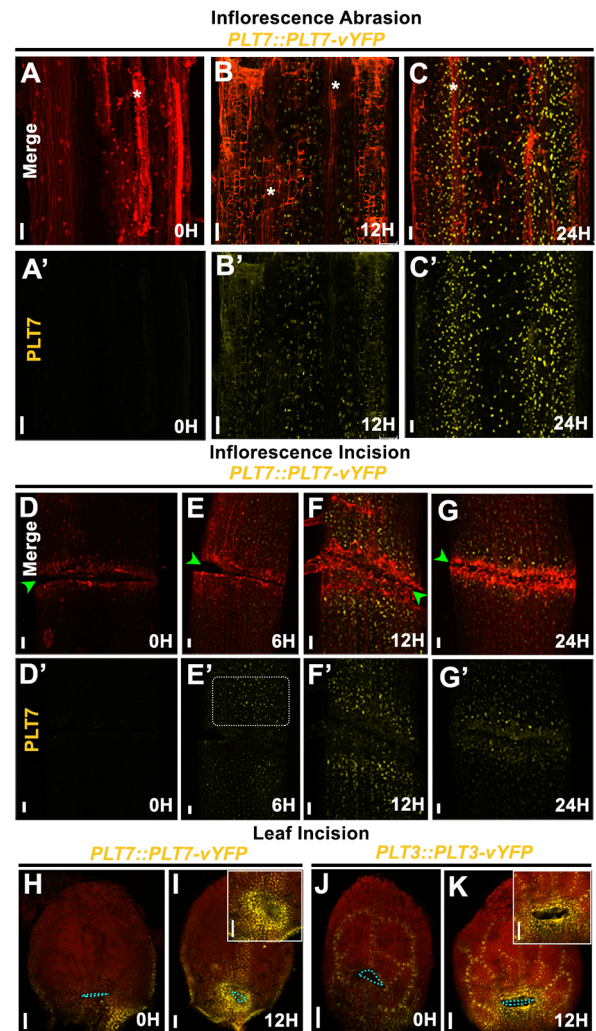
## RESULTS

### *PLT3*, *PLT5* and *PLT7* genes respond dynamically to mechanical injuries

*PLT3*, *PLT5* and *PLT7* collectively regulate tissue culture-mediated *in vitro* shoot regeneration and will from here on be referred to as *PLT3,5,7*. *PLT3,5,7*-regulated root stem cell regulators establish pluripotency in calluses and *PLT3,5,7*-regulated shoot-promoting factors act in response to external hormonal cues to induce regeneration of the complete plant body (Kareem et al., 2015).

Interestingly, *PLT3,5,7* genes are expressed in the shoot during development and positioning of aerial organs (Prasad et al., 2011; Krizek, 2015). To assess whether *PLT3,5,7* function is required for repairing damaged inflorescence and leaf tissue without external hormonal cues, we determined whether expression of these genes is induced as a natural response to injuries that growing plants are likely to encounter, such as local abrasions in the stem, partial stem incisions and midvein injuries in the leaf blade. These injuries were made without detaching any organ from growing *Arabidopsis* plants. After local abrasion that damaged the epidermal and sub-epidermal layers, including vascular tissue, in inflorescence stem (Fig. 1A,A', Fig. S1A-D), *PLT7::PLT7-vYFP* was induced 12 h post-injury, prior to any apparent regeneration response (Fig. 1B,B'). The expression peaked at 36 h (Fig. 1C,C', Fig. S1F,F'). In response to partial incision of the inflorescence stem (Fig. 1D,D'), *PLT7::PLT7-vYFP* expression was upregulated at both ends of the incised stem, with relatively higher expression in the upper end after 6 h (Fig. 1E,E'). The high level of expression continued for 12 h (Fig. 1F,F'). At 12 h, upregulated expression expanded beyond the partial slit, and at 24 h it became confined to a narrower domain in the vicinity of the incision (Fig. 1G-G'). Transcript levels of *PLT7* were consistent with the fusion protein expression data and remained upregulated at 24 h (Fig. S1E). Similarly, when the midvein of a growing leaf blade was wounded, cells in the vicinity displayed pronounced upregulation of *PLT7::PLT7-vYFP* 12 h post-injury (Fig. 1H,I). In response to injury, *PLT3::PLT3-vYFP* and *PLT5::PLT5-vYFP* also showed upregulation of expression in the vicinity of the wound albeit with some differences in the timing of their activation and in spatial distribution (Fig. 1J,K, Figs S1G-P', S2 and S3). In response to leaf incision, although both *PLT3* and *PLT7* were expressed in close proximity to the wound, *PLT5* was expressed predominantly in the vascular tissue near the damage (Fig. 1H-K, Fig. S3A,B).

The root stem cell regulators *PLT1*, *PLT2* and *WOX5*, which are activated by *PLT3,5,7* during tissue culture-mediated *in vitro* shoot regeneration (Kareem et al., 2015), were not expressed in growing leaves and stems in response to injuries (Fig. S4).



**Fig. 1. *PLT3*, *PLT5* and *PLT7* genes are locally induced after mechanical injury.** (A-G') *PLT7::PLT7-vYFP* expression (yellow) after abrasion (A-C') and partial incision (green arrowheads) (D-G') in growing inflorescence stems. White asterisks indicate vascular tissues exposed by damage to epidermal and sub-epidermal layers following local abrasion. E' white dotted area highlights upregulation of *PLT7* expression at the upper end of the cut. A'-C' and D'-G' are maximum intensity projections of z-stacks in the YFP channel corresponding to the regions shown in A-C and D-G, respectively. (H-K) Upregulation of *PLT7::PLT7-vYFP* (H,I) and *PLT3::PLT3-vYFP* (J,K) (yellow) near wound site (insets) following leaf incision (blue dotted area indicates the incision site). The panels represent different samples at each time point. Red signal is propidium iodide staining in A-G and chlorophyll autofluorescence in H-K. Brightness of the YFP signal was increased for visibility in B, B' and E'. H, hours after injury. Scale bars: 50  $\mu$ m.

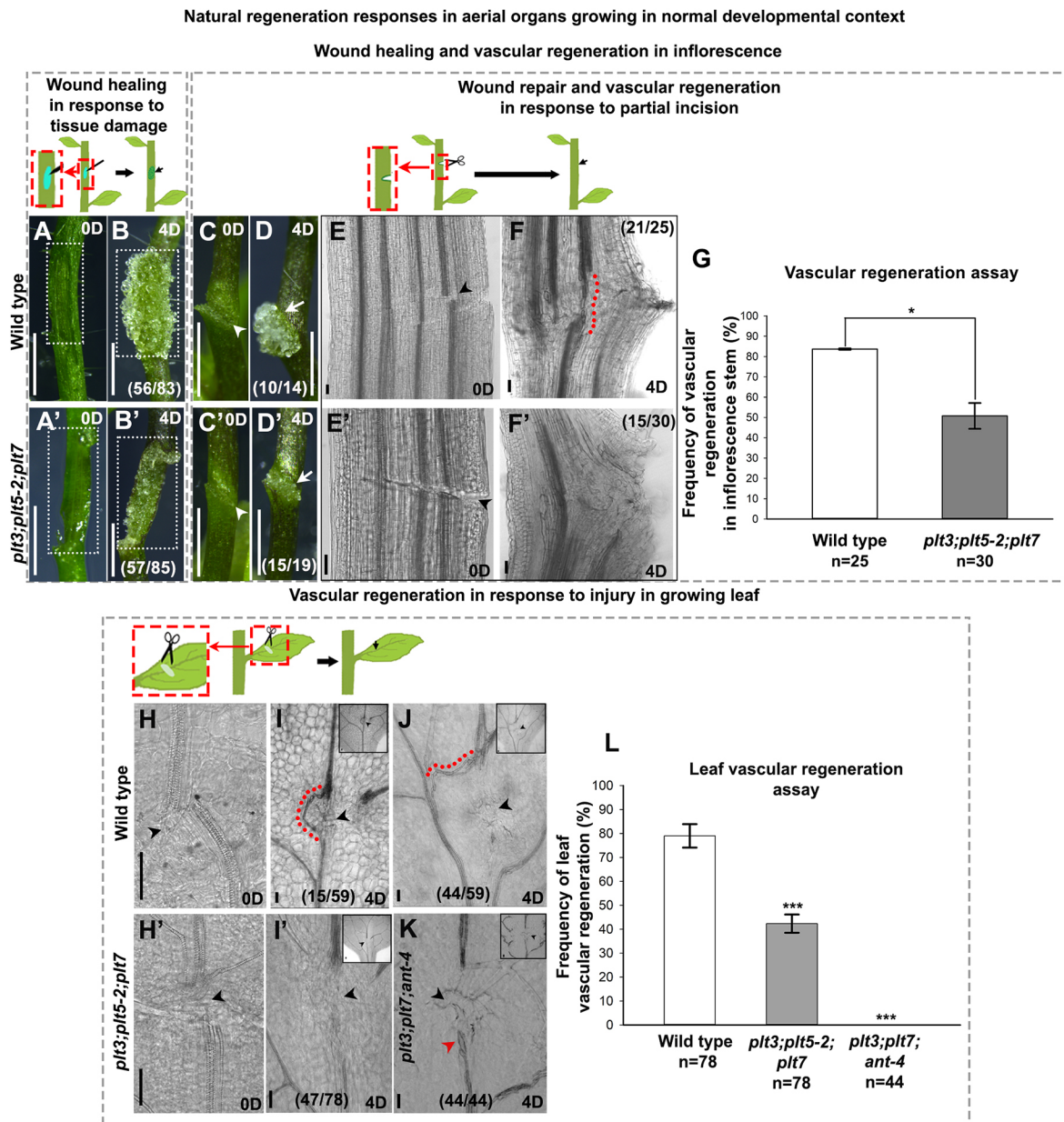
### *PLT3,5,7* are required to activate innate regenerative responses to injuries in aerial organs

Aerial organs of growing plants are subject to substantial wear and tear and *PLT3,5,7* expression is rapidly activated in response to injuries (Fig. 1, Figs S1-S3). We therefore asked whether *PLT3,5,7* genes are required for wound repair and tissue regeneration in stems and leaves growing in the normal developmental context of *Arabidopsis*.

#### Wound repair and vascular regeneration in inflorescence stem

We mimicked physical abrasion by damaging the epidermis, sub-epidermal layers and vascular tissue locally (see Materials and Methods for details; Fig. 2A,A') in a growing inflorescence stem of wild-type as well as *plt3;plt5-2;plt7* mutant plants. In the wild type,





**Fig. 2. *PLT* genes activate innate regenerative responses to injuries in aerial organs growing in the normal developmental context.** (A-F') Schematics above A and B represent inflorescence stem abrasion (red rectangle; cyan indicates wounded region) and wound healing response (arrow). Wound healing and vascular regeneration in inflorescence stems. (A,A') Abrasion (dotted rectangles) in inflorescence stems of wild type (A) and *plt3;plt5-2;plt7* (A'). (B,B') Reduced wound healing response (dotted rectangles indicate area of cell proliferation) in *plt3;plt5-2;plt7* (B') compared with wild type (B). (C,C') Partial incision (white arrowheads) in inflorescence stems of wild type (C) and *plt3;plt5-2;plt7* (C'). (D,D') Compromised callus formation (white arrows) in inflorescence stems of *plt3;plt5-2;plt7* (D') compared with wild type (D). (E,E') Disruption of vascular tissue (black arrowheads) by partial incision in inflorescence stems of wild type (E) and *plt3;plt5-2;plt7* (E'). (F,F') Vascular strands regenerate in wild-type inflorescence stems (F) but fail to regenerate in ~49% of *plt3;plt5-2;plt7* stems (F'). Schematics above E and F indicate partial incision on inflorescence stem (red rectangle) and wound healing response. Black arrow indicates site of wound healing. (G) Frequency of vascular regeneration in response to partial incision in the inflorescence stems of wild type and *plt3;plt5-2;plt7* (\* $P=0.033$ ; Pearson's  $\chi^2$  test). Schematics above H and I indicate incision in the midvein of the growing leaf (red rectangle). The wound is repaired by vascular regeneration and local cell proliferation. (H-K) Vascular strand regeneration in the growing leaf. (H,H') Incision (black arrowheads) in the midvein of wild-type (H) and *plt3;plt5-2;plt7* (H') growing leaves. (I) Vascular strands regenerate in the wild-type leaf, bypassing the wounded area and connecting the cut ends of the midvein. (J) A new vascular strand connects the upper cut end of the midvein to the lateral vein. Red dotted line in I and J indicate the regenerated vascular strand. (I',K) Vascular strands failed to regenerate in 60% of *plt3;plt5-2;plt7* leaves (I'). *plt3;plt7;ant-4* (K) mutant leaves completely failed to regenerate in response to midvein injury. Red arrowhead indicates proliferating cells at the lower cut end of the midvein. Insets show lower magnification images of the site of injury. Black arrowheads indicate the incision site. (L) Frequency of leaf vascular regeneration in wild type, *plt3;plt5-2;plt7* mutants (\*\* $P=1.211 \times 10^{-15}$ ; Pearson's  $\chi^2$  test) and *plt3;plt7;ant-4* mutants (\*\* $P=7.707 \times 10^{-13}$ ; Pearson's  $\chi^2$  test). Error bars represent s.e.m. In image panels, sample numbers are shown in parentheses. Scale bars: 1 mm (A-D'); 50  $\mu$ m (E-F', H-K). D, days after injury.

we noticed a healing response in the form of a visible mass of proliferating cells (callus-like growth) throughout the wound at 2 days after abrasion (daa), which became more prominent at 4 daa

(Fig. 2B, Fig. S5A,B). Later, callus-like growth completely covered and sealed the wound. The inflorescence stem regained its growth following the repair process. In contrast to wild type, the healing



response was severely reduced in injured *plt3;plt5-2;plt7* inflorescence stems and the wound-sealing process was not completed in the triple mutant inflorescence stem (Fig. 2A',B', Fig. S5A',B'). Importantly, the inflorescence stem development of uninjured mutant was comparable to that of wild type (Fig. S5I,J).

Next, we made a partial slit in the inflorescence stem of wild-type and *plt3;plt5-2;plt7* mutant plants disrupting both vascular connections and ground tissue (Fig. 2C,C',E,E'). Twenty-four hours after the incision, the wounded parts adhered in the wild-type inflorescence stem (Fig. S5C,D). Subsequently, cell proliferation was observed as indicated by visibly swollen tissues at cut ends followed by regeneration of vascular tissues at 4 days after cut (dac) (Fig. 2D,F). Subsequent restoration of growth and physiological functions were demonstrated by the development of new flowers and siliques (Fig. S5E). In contrast to wild type, in which the wound was healed on the fourth day, the *plt3;plt5-2;plt7* triple mutant displayed severely reduced callus-like growth at the wound site and ~49% inflorescence stems failed to regenerate vascular tissue (Fig. 2C'-F', G, Fig. S5C'). Our data demonstrate the role of *PLT3,5,7* in activating a healing response in the form of callus-like growth and vascular regeneration to restore damaged tissue in a growing inflorescence stem.

### Vascular regeneration in a growing leaf

Restoration of vasculature is a long-known feature of stem regeneration, and we investigated whether this response also occurred in leaves. We made a local injury in the midvein of a growing young wild-type leaf of a 5 days post-germination (dpg) plant (see Materials and Methods). To keep the developmental stage uniform, we injured the first pair of young leaves, which displayed midvein formation but not fully developed lateral veins at the time of injury (Fig. 2H,H', Fig. S6A,A'). The injuries either (1) damaged the midvein without making an opening or (2) completely disconnected the midvein leaving a gap between the vascular strands. In both the cases, cells in the vicinity of the midvein experienced mechanical perturbations due to the pressure applied by the needle. Four days post-injury (dpi), wild-type leaves repaired both types of injuries. In case (1), where the break was incomplete, the injured midvein was repaired and new vascular cells regenerated to restore the physiological connection (Fig. S6E). In case (2), where there was a complete disconnection, we observed regeneration of the vascular strand either connecting together the cut ends of the midvein or connecting the cut end of the midvein to a lateral vein (Fig. 2I,J, Fig. S6F,G). Strikingly, after local injury in the midvein of young wild-type leaf blades, ~80% of the samples regenerated vascular tissue in response to incision (Fig. 2L). The regenerating vascular cells often bypassed the damaged area and reunited with the lower half of the midvein making a D-shaped loop around the wound site similar to Sachs' observation of vascular regeneration around the wound site in the epicotyl stem of pea plants (Sachs, 1981) (Fig. 2I). Alternatively, they formed a new connection to the nearest lateral vein (Fig. 2J). The non-regenerating lower vascular strand degenerated after residual proliferation at the cut end (Figs S5L and S6B). We followed vascular regeneration from the time of injury to distinguish between the vascular strand reuniting the midvein regenerating from the cut end and the recruitment of a pre-existing lateral vein developed during leaf growth (Fig. S6A-D'). When the injury left a hole in the leaf blade exceeding 400 µm between the cut ends of the midvein, we rarely observed any vascular regeneration (Figs S6H,I and S7A). Such injuries left behind only a disorganised mass of cells (Fig. S6H). We therefore restricted our subsequent analysis to leaf blade injuries that completely disconnected the midvein leaving a gap well under 400 µm between the cut ends.

In contrast to wild-type leaves, in which ~80% of the injured leaves regenerated vascular strands, only ~40% of injured *plt3;plt5-2;plt7* leaves could regenerate and the rest completely failed to regenerate vascular strands (Fig. 2H',I',L). In non-regenerating mutant leaves, lateral veins failed to connect to the midvein near the wound site (Fig. 2I'), but a disorganised mass of proliferating cells at the wound site was observed, mostly at the cut ends of upper vascular strands and on the epidermis (Fig. S7B-D). Many such leaves displayed poor growth and failed to develop properly (Fig. S7E). It is important to note that uninjured *plt3;plt5-2;plt7* mutant plants did not display any defects in the formation of closed vein loops (consisting of midvein and secondary veins) compared with wild type but were severely impaired in vascular regeneration (Fig. 2L, Fig. S7F-I). With respect to leaf morphology, we did not observe any defects in the first pair of leaves (Fig. S5F,G). Among double mutant combinations, 70% of *plt3;plt5-2* and *plt5-2;plt7* double mutants regenerated vascular strands in response to injury, and only ~64% of *plt3;plt7* double mutant leaves regenerated vascular tissue (Fig. S8A).

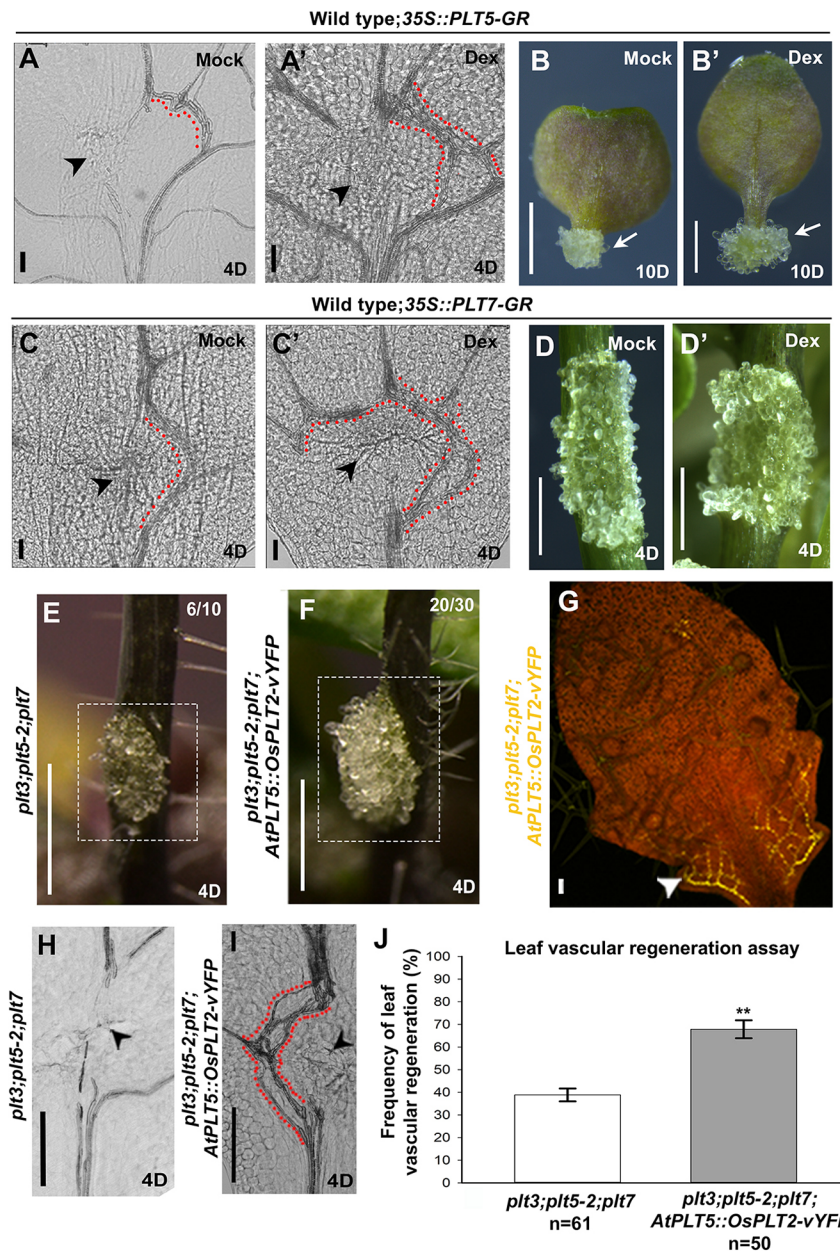
The closely related *AJINTEGUMENTA* (*ANT*) gene marks stem cells of root vascular cambium and acts redundantly with *PLT3* and *PLT7* during plant development (Krizek, 2015; Smetana et al., 2019). *ANT* is strongly expressed in the vascular tissue of young leaves (Fig. S8B). We therefore examined vascular regeneration in *plt3;plt7;ant-4* triple mutant plants in response to midvein injury. Strikingly, none of the tested *plt3;plt7;ant-4* seedlings regenerated vascular tissues, demonstrating an essential role of *ANT* with *PLT3* and *PLT7* in vascular regeneration (Fig. 2K,L, Fig. S8C). Taken together, our data reveal a previously unrecognised role of *PLT3,5,7* and *ANT* in repairing damaged tissues during plant growth.

Because of the severity of shoot phenotypes in *plt3;plt7;ant-4* (which produces only leaves but no stem) we chose the *plt3;plt5-2;plt7* mutant, which develops normal leaves as well as an inflorescence stem comparable to wild type, to probe the mechanism of vascular regeneration using further assays (Prasad et al., 2011; Krizek, 2015).

### *PLT5* and *PLT7* are sufficient for promoting vascular regeneration and wound repair

Tissue/organ regeneration is closely linked to cellular reprogramming. We next asked whether *PLT* genes are sufficient to activate cellular reprogramming leading to enhancement of wound repair. Strikingly, inducible overexpression of *PLT5* (*35S::PLT5-GR*) or *PLT7* (*35S::PLT7-GR*) promoted multiple strand formation from the regenerating midvein in response to injury (Fig. 3A,A',C,C', Fig. S8D). Similarly, inducible overexpression of *PLT5* or *PLT7* enhanced wound repair at the cut ends of the detached organ and in response to inflorescence abrasion (Fig. 3B,B',D,D', Fig. S8E-F'). Consistent with the ability of *PLT* proteins to promote cell division upon wounding, transcripts of *CYCLIN* genes increased in growing seedlings upon inducible overexpression of *PLT5* (*35S::PLT5-GR*) (Fig. S8G). These results suggest that *PLT* proteins are sufficient to promote wound repair and multiple vascular strand regeneration in response to injury.

We addressed whether *PLT*-like proteins from other plant species can trigger regeneration in *Arabidopsis*. Rice is a morphologically diversified monocot plant, whereas *Arabidopsis* is a dicot. Expression of the rice *PLT*-like gene *OsPLT2* under the *Arabidopsis* *PLT5* promoter in a *plt3;plt5-2;plt7* mutant (*plt3;plt5-2;plt7;AtPLT5::OsPLT2-vYFP*) healed a damaged *Arabidopsis* *plt* mutant inflorescence stem by inducing cell proliferation as evident from upregulated expression of cell cycle progression markers (Fig. 3E-G, Fig. S8H,I). Furthermore, *OsPLT2-vYFP* rescued leaf vascular regeneration defects in *plt3;plt5-2;plt7* suggesting that it is a functional homologue of *Arabidopsis* *PLT* genes (Fig. 3H-J).



**Fig. 3. *PLT* genes are sufficient for enhancing vascular regeneration and wound repair.**

(A-B') Overexpression of 35S::*PLT5-GR* promotes multiple vascular strand (A') formation upon leaf incision and callus formation (white arrows) at cut end of detached organ (B') unlike in mock-treated control (A,B). (C-D') Overexpression of 35S::*PLT7-GR* enhances multiple strand formation upon leaf incision (C') and wound repair upon inflorescence abrasion (D') unlike in mock-treated control (C,D). (E,F) Only a residual cell proliferation response is observed in *plt3;plt5-2;plt7* (E) unlike the extensive callus-like growth observed in *plt3;plt5-2;plt7;AtPLT5::OsPLT2-vYFP* (F) in response to inflorescence abrasion. Dotted rectangle indicates area of cell proliferation. (G) Expression of *AtPLT5::OsPLT2-vYFP* in vascular tissue (white arrowhead) of a *plt3;plt5-2;plt7* leaf. (H-J) Rescue of vascular tissue regeneration in response to leaf incision in *plt3;plt5-2;plt7;AtPLT5::OsPLT2-vYFP* (I,J) (\*\**P*=0.004; Pearson's  $\chi^2$  test) compared with *plt3;plt5-2;plt7* leaves (H), of which ~61% failed to regenerate. Error bars represent s.e.m. Black arrowheads indicate incision site. Red dotted lines indicate regenerated vascular strands. Dex, dexamethasone. Scale bars: 50  $\mu$ m (A,A',C,C',G-I); 1 mm (B,B',D-F). D, days after injury.

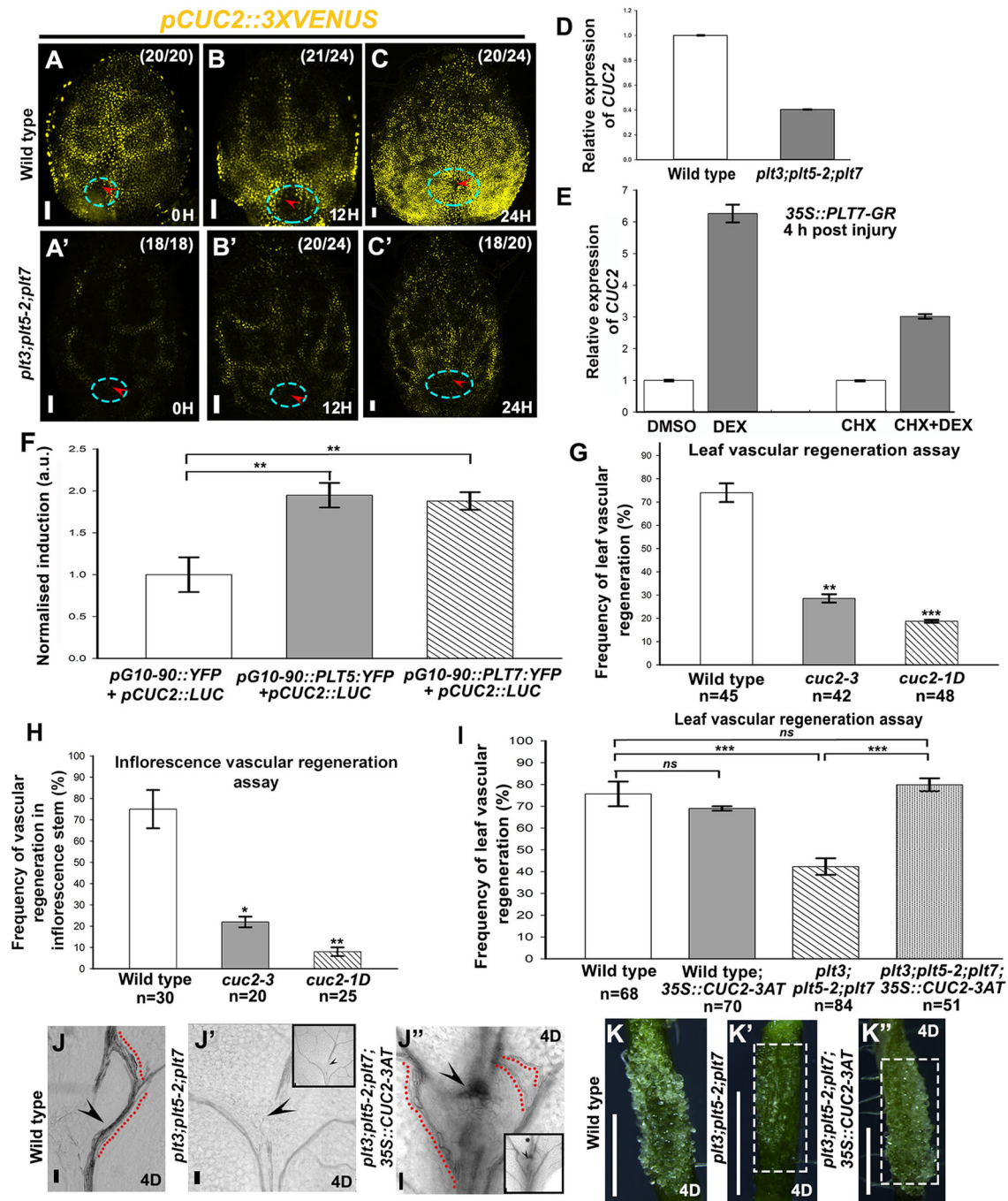
### ***PLT3,5,7* directly activate *CUC2* expression for wound repair and vascular regeneration**

Having established that *PLT3,5,7* regulate wound repair and vascular regeneration in damaged aerial parts of the plant, we sought to define the molecular mechanisms underlying this regulation. Previously, we had shown that *PLT3,5,7* direct tissue-culture-mediated *in vitro* shoot regeneration by activating root stem cell regulators and *CUC2* (Kareem et al., 2015). Although we found no evidence for the participation of *PLT1*, *PLT2* and *WOX5* root stem cell regulators in the response to injuries in growing aerial organs (Figs S4 and S9A), *CUC2* remains an attractive candidate for participation in wound repair. Therefore, we investigated whether *CUC2* responds to mechanical injury and whether *PLT3,5,7* act through *CUC2* to repair wounds and regenerate vasculature.

*pCUC2::3XVENUS* as well as *CUC2::CUC2-vYFP* expression was detected in vascular tissue of young leaves in both wild-type and *plt3;plt5-2;plt7* mutant plants although expression was reduced in the latter (Fig. 4A,A', Fig. S9B-F,I,I'). The same *CUC2* promoter was

used to drive transcriptional and translational fusions. The detection of an expanded domain of expression of *pCUC2::3XVENUS* compared with *CUC2::CUC2-vYFP* can be largely attributed to *3XVENUS*. Both reporter fusions used in this study can recapitulate the previously reported *CUC2* expression at the leaf margin (Nikovic et al., 2006; Bilsborough et al., 2011) (Fig. S9G,H). In response to midvein damage in wild type, expression of both *pCUC2::3XVENUS* and *CUC2::CUC2-vYFP* was upregulated proximal to the wound 12 h post-injury followed by a broader domain of enhanced expression after 24 h (Fig. 4B,C, Fig. S9I-K). In contrast, there was no upregulation of the reporter near the wound site in *plt3;plt5-2;plt7* (Fig. 4B',C', Fig. S9I'-K'). Similar patterns of changes were also observed at the transcript level in response to midvein injury (12 h post-injury) (Fig. S9L). Similarly, in damaged inflorescence stems, *CUC2* transcripts were reduced in the *plt3;plt5-2;plt7* compared with wild type (Fig. 4D). Furthermore, *CUC2* transcripts rapidly increased in injured leaves upon inducible overexpression of *PLT5* (35S::*PLT5-GR*) as well as of *PLT7* (35S::*PLT7-GR*) even in the presence of the





**Fig. 4. *PLT* genes act through *CUC2* to repair wounds and to regenerate vascular tissue.** (A–C') Reduced expression of *pCUC2::3XVENUS* (yellow) in *plt3;plt5-2;plt7* (A'–C') compared with wild type (A–C) in response to injury. Red arrowheads denote incision site and dashed circles enclose leaf tissue in the vicinity of the wound showing upregulation of *pCUC2::3XVENUS* in wild type but not in *plt3;plt5-2;plt7*. Sample numbers are shown in parentheses (numerator, number of samples showing the expression represented in the image panel; denominator, total number of samples analysed). (D) Relative expression levels (qRT-PCR) of *CUC2* in injured *plt3;plt5-2;plt7* mutant inflorescence segments compared with wild type (4 dpi). (E) Rapid upregulation of *CUC2* (qRT-PCR) in injured tissue upon induction of 35S::PLT7-GR. Expression levels in D and E are normalised to *ACTIN2*. Error bars represent s.e.m. from three independent biological replicates. (F) PLT5 and PLT7 induce *pCUC2* in a luciferase reporter assay 2 days post-inoculation in *Nicotiana*. \*\**P*<0.01 (Mann–Whitney *U* one-tailed test). Six biological replicates each with three technical replicates were performed. (G) Frequency of leaf vascular regeneration in *cuc2-3* (recessive) (\*\**P*=0.007), *cuc2-1D* (dominant) (\*\**P*=0.0005) mutants compared with wild type (Pearson's  $\chi^2$  test). (H) Frequency of vascular regeneration in response to partial incision in the inflorescence stem of wild type, *cuc2-3* and *cuc2-1D* (\**P*=0.02, \*\**P*=0.001; Pearson's  $\chi^2$  test). (I) Frequency of leaf vascular regeneration in wild type, wild type; 35S::CUC2-3AT (ns, not significant; *P*=0.65), *plt3;plt5-2;plt7* (\*\**P*=9.9×10<sup>−5</sup>) and *plt3;plt5-2;plt7*; 35S::CUC2-3AT (\*\**P*=4.7×10<sup>−6</sup>) (Pearson's  $\chi^2$  test). (J) Representative example of vascular regeneration in a wild-type leaf. (J', J'') Vascular tissue regeneration is rescued in *plt3;plt5-2;plt7*; 35S::CUC2-3AT (J'') compared with *plt3;plt5-2;plt7* (J') in response to leaf incision (black arrowheads). Note the increased vascular strand proliferation and regeneration of multiple vascular strands (red dotted lines) generating multiple reunion points in *plt3;plt5-2;plt7*; 35S::CUC2-3AT (J'') unlike in *plt3;plt5-2;plt7* (J'). Insets show the incision site. (K) Representative example of local cell proliferation response in a wild-type inflorescence stem in response to abrasion. (K', K'') Ectopic overexpression of *CUC2* in *plt3;plt5-2;plt7* (*plt3;plt5-2;plt7*; 35S::CUC2-3AT) (K'') enhances local cell proliferation and wound healing response upon inflorescence abrasion (enclosed by dashed rectangle) compared with *plt3;plt5-2;plt7* (K'). Error bars represent s.e.m. Scale bars: 1 mm (K–K''); 50  $\mu$ m (A–C', J–J''). D, days after injury; H, hours after injury.



translation inhibitor cycloheximide, suggesting direct activation of *CUC2* transcription by PLT5 and PLT7 (Fig. 4E, Fig. S9M). Consistent with these observations, PLT5 bound to the *CUC2* promoter in a chromatin immunoprecipitation (ChIP) assay (Fig. S9N). In addition, DNA affinity purification sequencing (DAP-Seq) analysis identified the binding of PLT7 to the *CUC2* promoter (O'Malley et al., 2016, <http://neomorph.salk.edu/>) (Fig. S10A). Furthermore, transient transfection of trans genes capable of producing PLT5 or PLT7 proteins and the *CUC2* promoter-driven luciferase reporter in *Nicotiana* leaf induced reporter expression, further demonstrating that PLT5 as well as PLT7 can directly activate *CUC2* transcription (Fig. 4F).

Because molecular data indicate that *CUC2* acts downstream of *PLT* genes, we investigated whether PLT proteins require *CUC2* activity for wound repair. Strikingly, inducible ectopic overexpression of PLT5 failed to promote wound repair at the damaged end of *cuc2-3* (*cuc2-3;35S::PLT5-GR*) mutant tissues. The severely compromised wound repair that was observed at the cut ends remained unaltered upon PLT5 overexpression in *cuc2-3* detached tissue, but not upon PLT5 overexpression in wild type (Wild type;*35S::PLT5-GR*), which enhanced wound repair at the cut ends (Fig. S10B-F). These results demonstrate that PLT proteins act through *CUC2* to repair the wound.

We examined the role of *CUC2* in leaf vascular regeneration by analysing loss-of-function mutants. Strikingly, vascular regeneration was severely impaired in both the recessive loss-of-function *cuc2-3* mutant as well as in the *cuc2-1D* dominant mutant; 71% of *cuc2-3* mutant and 81% of *cuc2-1D* mutant leaves failed to show any vascular regeneration in response to midvein injury (Fig. 4G). Notably, loss of *CUC2* function did not cause any defect in the formation of closed vein loops formed by primary (midvein) and secondary (lateral) veins (Fig. S7F,G,J,K,L). Similarly, upon inflorescence stem incision, ~78% *cuc2-3* mutant and 92% of *cuc2-1D* mutant inflorescence stems failed to show any vascular regeneration (Fig. 4H). Finally, we asked whether *CUC2* overexpression can rescue the vascular regeneration defect in *plt3;plt5-2;plt7* mutant leaves. Strikingly, the regeneration efficiency (timings of regeneration and reunion of vascular strands) as well as frequency (number of plants) was restored upon *CUC2* overexpression in *plt3;plt5-2;plt7* to the level of wild type. New vascular strands regenerated and reunited by 4 dpi in the mutant, similar to wild type (Fig. 4I-J"). Moreover, *CUC2* overexpression rescued the repair process in locally wounded *plt3;plt5-2;plt7* inflorescence stems (Fig. 4K-K"). Taken together, these data demonstrate that *PLT3,5,7* directly activate *CUC2* transcription in response to injury and that the *PLT-CUC2* module is required for wound repair and vascular regeneration in leaf and stem. Interestingly, the growth of inflorescence stems and leaves were similar in wild type and the *cuc2-3* mutant (Fig. S5H,K).

### **PLT genes are required for polarised cell growth and auxin response during vascular regeneration**

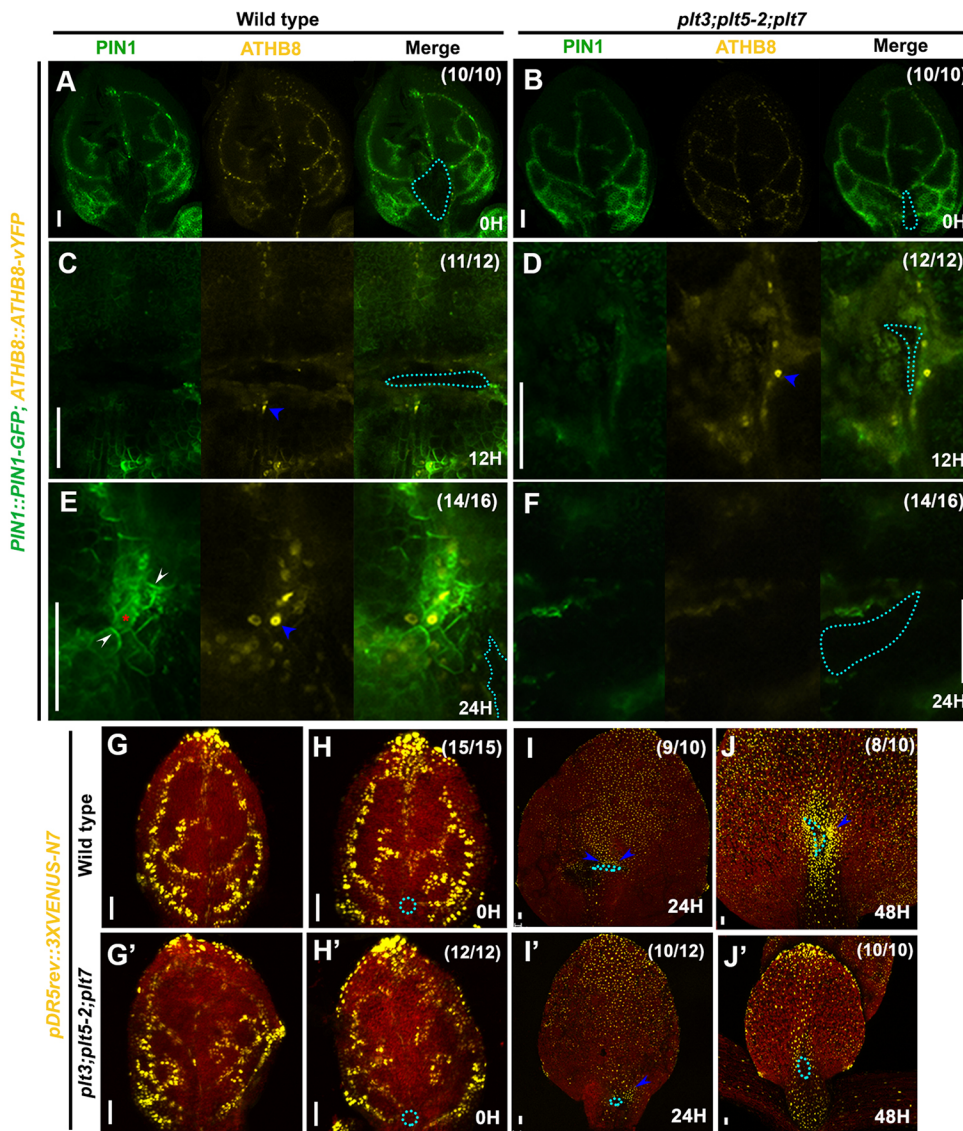
*CUC2* is implicated in the regulation of leaf margin development by directing PIN1 polarity and the resultant auxin distribution (Bilborough et al., 2011) and PIN1 polarisation is crucial for the normal development of leaf vasculature (Scarpella et al., 2006). Hence, we next investigated whether the process of cell polarisation is regulated by the *PLT* transcription module during leaf vascular regeneration. We focused on *in vivo* vascular regeneration in developing leaves, which has not been explored. To this end, we examined the localisation of PIN1 in response to midvein injury in the leaf blade (Fig. S11C'-F',G'-J'). Prior to wounding, we observed

*PIN1::PIN1-GFP* expression in the procambium predominantly towards the basal end of young leaves in both wild type and mutant (Fig. S11A-F,G-J). In response to injury, we observed increased PIN1-GFP near wound sites in both wild type and mutant (Fig. S11E',I'). To examine PIN1-GFP localisation in regenerating vascular cells, we generated transgenic lines harbouring both *PIN1::PIN1-GFP* and *ATHB8::ATHB8-vYFP*. *ATHB8* specifically marks developing procambium cells in leaf (Scarpella et al., 2004). We observed both PIN1-GFP and *ATHB8-YFP* in developing procambium of 4-day-old leaves (Fig. 5A,B).

During the first 12 h following incision, we did not observe regenerating vascular cells expressing both PIN1-GFP and *ATHB8-YFP* near wound sites (Fig. 5C,D). Regenerating procambium cells marked with *ATHB8-YFP* and polarised PIN1-GFP were observed after 24 h near wound in wild type (Fig. 5E). In contrast, we did not observe regenerating procambium cells expressing polarised PIN1-GFP or *ATHB8-YFP* near the wound after 24 h in *plt3;plt5-2;plt7* plants, demonstrating that cells surrounding the damaged site failed to re-specify the PIN1 polarity in the mutant (Fig. 5F). These data suggest that failure of re-establishment of polar auxin transport within 24 h may contribute to impaired vascular regeneration in the *plt* triple mutant. We next examined whether lack of directional auxin flow in the damaged *plt3;plt5-2;plt7* mutant leaves altered the distribution patterns of the auxin response. We examined the auxin response using the auxin reporter *pDR5rev::3XVENUS-N7* in both wild-type and *plt3;plt5-2;plt7* mutant plants. Prior to injury, we did not observe any difference in distribution patterns or levels of the auxin response in leaves between these two genotypes (Fig. 5G,G', Fig. S11K-M'). In both wild type and *plt3;plt5-2;plt7* an increase in *pDR5rev::3XVENUS-N7* signal in the tissue proximal to the wound was observed at 24 h post-injury (Fig. 5H,H',I,I'). However, a further enhanced auxin response was confined to an area near the wound site by 48 h only in wild type (Fig. 5J). In contrast to wild type, the triple mutant failed to show such confined expression of *pDR5rev::3XVENUS-N7* signal in response to injury (Fig. 5J'). The distribution patterns and levels of auxin response in uninjured developing mutant leaves compared with wild type did not change, further substantiating the specific role of *PLT3,5,7* in response to injury (Fig. 5G,G', Fig. S11K-M'). Taken together, our results show that *PLT3,5,7* are needed for re-specification of polarised vascular cells to facilitate vascular tissue regeneration.

### **PLT proteins and CUC2 activate the transcription of local auxin biosynthesis gene in a feed-forward loop to repair wounds and drive vascular regeneration**

Local auxin biosynthesis has been implicated in root haustoria formation and associated vascular development during host-parasite interaction (Kokla and Melnyk, 2018). We therefore asked whether *PLT-CUC2* module regulate wound repair and vascular regeneration by modulating local auxin biosynthesis genes. Interestingly, local auxin biosynthesis genes are downregulated in *plt3;plt5-2;plt7* mutant callus (A.K. and K.P., unpublished data). PLT proteins are also known to control phyllotaxis by regulating one of the auxin biosynthesis genes, *YUCCA4* (*YUC4*) (Pinon et al., 2013). Similarly, *YUC4* expression was upregulated in response to midvein injury (12 h post-injury) in growing wild-type leaves, unlike in the *plt3;plt5-2;plt7* leaves, in which the transcript level was reduced (Fig. 6A). In addition to damaged leaves, *YUC4* transcripts were also reduced in damaged *plt3;plt5-2;plt7* inflorescence segments (Fig. S12A). Conversely, *YUC4* transcripts were rapidly increased in injured tissues upon PLT5-GR induction (4 h) even in the presence of the translation inhibitor cycloheximide, suggesting direct activation by



**Fig. 5. *PLT* genes regulate polarised cell growth and auxin response during vascular regeneration.** (A-F) Expression of *PIN1::PIN1-GFP* and *ATHB8::ATHB8-vYFP* in wild-type (A,C,E) and *plt3;plt5-2;plt7* mutant (B,D,F) leaves in response to leaf incision. (A,B) YFP channel shows expression of *ATHB8::ATHB8-vYFP* in procambium cells of the leaf. (C,D) No expression of *PIN1* is detected in the immediate vicinity of the wound at 12 h in either wild type (C) or *plt3;plt5-2;plt7* (D). Pre-existing *ATHB8* (blue arrowheads) expression is observed near the wound (in C and D YFP channel). (E,F) Expression of polarised *PIN1::PIN1-GFP* (white arrowheads) and *ATHB8::ATHB8-vYFP* (blue arrowhead) in the regenerating cells (hexagonal developing procambium cells; indicated by red asterisk) of wild type. *PIN1* polarisation and *ATHB8* expression is absent in *plt3;plt5-2;plt7* (F). Blue dotted area indicates tissue damaged by leaf incision. Brightness of YFP channel (representing *ATHB8*) signal was increased for visibility in A-F. A-F show a subset of z-stack sections. (G,G') *pDR5rev::3XVENUS-N7* expression in undamaged leaves of wild type (G) and *plt3;plt5-2;plt7* (G'). (H-J') *pDR5rev::3XVENUS-N7* expression in wild-type (H-J) and *plt3;plt5-2;plt7* (H'-J') leaves post-incision (dotted area indicates incision site). *plt* mutant leaf remains small due to stunted growth following injury. Note the confined expression of *pDR5rev::3XVENUS-N7* in the tissue around the wound (blue arrowheads) in wild type (J) unlike in *plt3;plt5-2;plt7* (J'). Scale bars: 50 μm. Red colour in G-J' represents chlorophyll autofluorescence. Sample numbers are shown in parentheses (numerator, number of samples showing the expression represented in the image panel; denominator, total number of samples analysed). H, hours after injury.

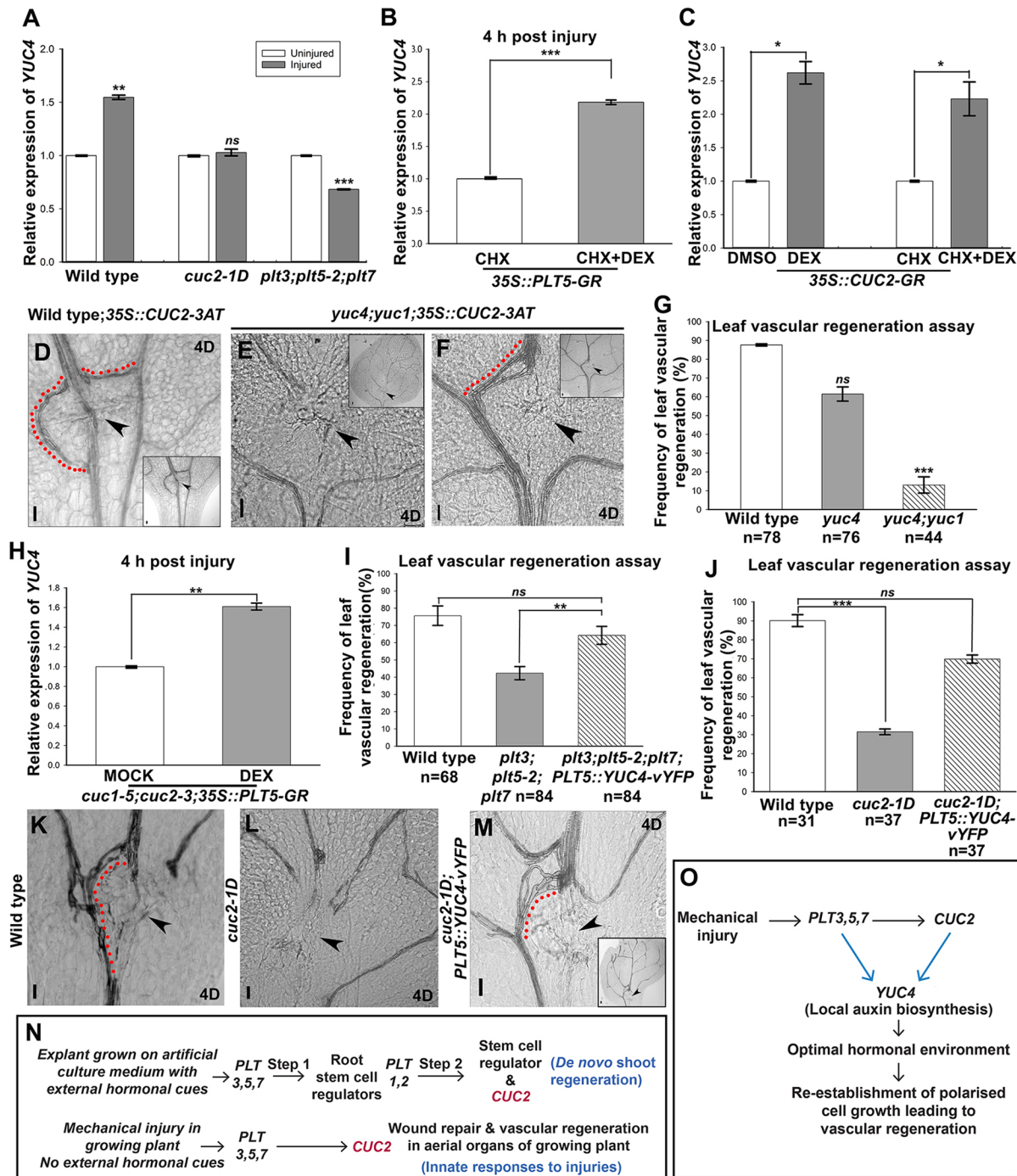
*PLT5* (Fig. 6B). Because molecular data suggests that *YUC4* acts downstream of *PLT* genes, we investigated whether *PLT* proteins require *YUC4* activity to trigger cellular reprogramming. Strikingly, inducible overexpression of *PLT5* as well as *PLT7* failed to trigger any ectopic cellular reprogramming in the *yuc4;yuc1* mutant background (*yuc4;yuc1;35S::PLT5-GR* or *yuc4;yuc1;35S::PLT7-GR*), unlike in the wild-type background (Wild type;*35S::PLT5-GR*; or Wild type;*35S::PLT7-GR*) (Fig. S12B,C). Similarly, *PLT5* as well as *PLT7* overexpression failed to promote wound repair at damaged ends, demonstrating that *PLT* proteins act through *YUC4* during reprogramming and wound repair (Fig. S12D-G).

We explored whether, in addition to *PLT* genes, *CUC2* might also contribute towards regulating the local auxin biosynthesis in response to injury. *YUC4* transcripts were not upregulated in response to midvein injury in the *cuc2-1D* single mutant (Fig. 6A) and its transcript levels were rapidly increased upon *CUC2-GR* induction even in the presence of the translation inhibitor cycloheximide, suggesting direct activation of *YUC4* expression by *CUC2* (Fig. 6C). Consistent with the likelihood of direct activation of *YUC4* transcription by *CUC2* inferred from our results, DAP-Seq analysis indicated the binding of *CUC2* to the *YUC4* promoter (Fig. S12H) (O'Malley et al., 2016, <http://neomorph.salk.edu/>). Next, we

examined whether, like *PLT* proteins, *CUC2* also requires downstream *YUC4* activity to promote vascular regeneration. Ectopic overexpression of *CUC2* promoted vascular regeneration in the leaf and resulted in regeneration of multiple vascular strands from the wound site in the wild type (Wild type;*35S::CUC2-3AT*) (Fig. 6D). In contrast to wild type, ectopic overexpression of *CUC2* failed to promote regeneration of multiple vascular strands from the wound site in the *yuc4;yuc1* mutant (*yuc4;yuc1;35S::CUC2-3AT*) (Fig. 6D-F). Injured leaves in *yuc4;yuc1;35S::CUC2-3AT* seedlings either did not regenerate any vascular strand or occasionally displayed a single file of regenerating vascular cells as was observed in *yuc4;yuc1* (Fig. 6E,F, Fig. S12I). These data demonstrate that, like *PLT* genes, *CUC2* also acts through *YUC4* to promote wound repair and vascular regeneration.

Our data suggest that, in addition to *PLT* proteins, *CUC2* can also activate *YUC4* expression during vascular regeneration. Activation of *YUC4* by *PLT5*, *PLT7* and *CUC2* indicates a feed-forward loop controlling local auxin biosynthesis (Fig. 6B,C, Fig. S12J). *PLT5-GR* can only moderately activate *YUC4* expression after 4 h induction when the function of *CUC2* and of the redundantly acting *CUC1* is lost (in damaged *cuc1-5;cuc2-3* tissues) (Fig. 6H), indicating that increased transcription of *YUC4* in wild-type damaged leaves may be





**Fig. 6. *PLT*- and *CUC2*-dependent auxin biosynthesis drives vascular regeneration in leaf.** (A) *YUC4* transcript level in wild-type, *cuc2-1D* and *plt3;plt5-2;plt7* injured and uninjured leaves, measured by qRT-PCR (ns, not significant,  $P=0.45$ ; \*\* $P=0.001$ , \*\*\* $P=0.0002$ ; Welch's two-sample *t*-test). (B) Upregulation of *YUC4* (qRT-PCR) transcript level in injured leaves upon induction of 35S::PLT5-GR with cycloheximide (CHX) treatment (\*\*\* $P=0.0008$ ; Welch's two-sample *t*-test). (C) Upregulation of *YUC4* (qRT-PCR) transcript levels in injured leaves upon induction of 35S::CUC2-GR with and without CHX treatment (\* $P<0.05$ ; Welch's two-sample *t*-test). (D-F) Ectopic overexpression of *CUC2* produced multiple vascular strands from the wound site in wild type;35S::CUC2-3AT (D) unlike in *yuc4;yuc1*;35S::CUC2-3AT (E,F). (G) Percentage of leaf vascular regeneration in wild type, *yuc4* (ns, not significant;  $P=0.8$ ) and *yuc4;yuc1* (\*\*\* $P=1.02\times10^{-6}$ ; Pearson's  $\chi^2$  test). (H) *YUC4* transcript level in *cuc1-5;cuc2-3* upon induction of 35S::PLT5-GR, measured by qRT-PCR. Data shown in A-C,H are normalised to *ACTIN2*. Error bars represent s.e.m. from three independent biological replicates (\*\* $P=0.0032$ ; Welch's two-sample *t*-test). (I) Frequency of leaf vascular regeneration in wild type, *plt3;plt5-2;plt7* (ns, not significant,  $P=0.18$ ) and *plt3;plt5-2;plt7;PLT5::YUC4-vYFP* (\*\* $P=0.0087$ , Pearson's  $\chi^2$  test). (J-M) Reconstitution of local auxin biosynthesis gene in the *PLT5* domain rescues leaf vascular regeneration in the *cuc2-1D* mutant (\*\*\* $P=4.11\times10^{-6}$ ; ns, not significant,  $P=0.08$ ). Error bars represent s.e.m. Black arrowheads indicate incision site. Red dotted lines indicate regenerated vascular strands. Insets in D-F,M show lower magnification images of the site of injury. Scale bars: 50  $\mu$ m. D, days after injury. (N) Schematic showing the *PLT*-*CUC2* module independently activating innate regeneration responses to injuries, which is unlike the sequential activation of *CUC2* after activation of root stem cell regulators during *de novo* shoot regeneration. (O) Schematic representing the mechanistic module of *PLT* transcription factors activating *CUC2* and *YUC4* to generate an optimal auxin environment to aid in re-establishment of polarised growth of vascular cells. Regulatory interactions marked using light blue arrows emerged from the present study and were not known previously in any regeneration context.



an output of a coherent feed-forward loop during tissue regeneration. We further provide genetic evidence for the feed-forward loop: inducible overexpression of *PLT7* or *PLT5* can still increase vascular regeneration by 18% and 24%, respectively, in response to midvein injury in the *cuc2-3* mutant (Fig. S12K).

We further investigated this regulatory interaction by analysing the genetic interaction between *PLT* genes and *CUC2*. Strikingly, we found synergistic interaction between *PLT* genes and *CUC2* during wound repair and vascular regeneration. Cumulative loss of *PLT* and *CUC2* function in the *plt3;plt5-2;plt7;cuc2-3* mutant resulted in severely compromised wound repair at the cut end of the detached plant organ compared with the *plt3;plt5-2;plt7* or *cuc2-3* mutant (Fig. S13A). In addition to dramatically reduced frequency of wound repair in *plt3;plt5-2;plt7;cuc2-3* mutant, we could barely observe any proliferating callus-like cells at the damaged ends in *plt3;plt5-2;plt7;cuc2-3* mutant organs (Fig. S13B-E). The *YUC4* transcript level was further reduced in the *plt3;plt5-2;plt7;cuc2-3* mutant compared with the *plt3;plt5-2;plt7* or *cuc2-3* mutant (Fig. S13F). Similarly, seedlings heterozygous for *plt* and *cuc2* alleles, *plt3<sup>+/-</sup>;plt5-2<sup>+/-</sup>;plt7<sup>+/-</sup>;cuc2-3<sup>+/-</sup>* displayed hypersensitivity to leaf midvein injury compared with *plt3<sup>+/-</sup>;plt5-2<sup>+/-</sup>;plt7<sup>+/-</sup>* or *cuc2-3<sup>+/-</sup>* (Table S1, Fig. S7L). These data substantiate the regulation of *YUC4* expression by *PLT* proteins and *CUC2* in a coherent feed-forward loop during wound repair and vascular regeneration.

Consistent with the importance of activation of *YUC4* expression for regeneration, ~40% of *yuc4* single mutant and 87% of *yuc4;yuc1* double mutant leaves failed to regenerate vascular tissue in response to midvein injury (Fig. 6G). Strikingly, the uninjured *yuc4* single mutant develops a fully grown midvein without any discontinuity and there is no significant difference in the formation of closed vein loops compared with wild type (Fig. S7F,G,M). Although midvein formation in the *yuc4;yuc1* mutant was normal, as in the wild type, the number of loops surrounding the midvein was reduced (Fig. S7F,G,N). Strikingly, reconstitution of *YUC4* expression in the endogenous *PLT5* domain (*PLT5::YUC4-vYFP*) in the *plt3;plt5-2;plt7* mutant as well as in the *cuc2-1D* mutant rescued the vascular regeneration in injured leaves to a large extent (Fig. 6I-M, Fig. S13G-I). These data provide compelling evidence for the functional significance of *PLT-CUC2* module-dependent activation of local auxin biosynthesis in controlling vascular regeneration. Remarkably, reconstitution of *YUC4* expression in the *cuc1-5;cuc2-3* (*cuc1-5;cuc2-3;PLT5::YUC4-vYFP*) mutant, which generates cup-shaped cotyledons but no leaf or stem, rescued post-embryonic development with fully developed rosette leaves (Fig. S14).

## DISCUSSION

Multicellular organisms display the ability to regrow damaged tissues and organs. Unlike many animals, in which regeneration potential is restricted to specific cell lineages, plants repair and rebuild damaged tissues throughout the body. In this study, we have investigated the mechanism of wound repair across aerial parts of the plant body, identifying *PLT/AIL* transcription factors, well known for their role in stem cell maintenance, as regulatory triggers for this process. We demonstrate that activation of *CUC2* transcription by *PLT3,5,7* is a key regulatory mechanism of wound repair and vascular regeneration: (1) *PLT* factors bind to the *CUC2* promoter and directly activate the transcription of *CUC2*; (2) *PLT* factors require downstream *CUC2* activity during wound repair; and (3) reconstitution of *CUC2* expression under a heterologous promoter in *plt3;plt5-2;plt7* triple mutants rescues vascular regeneration. We provide evidence that *PLT* proteins and

*CUC2* activate the transcription of local auxin biosynthesis gene in a feed-forward loop to drive vascular regeneration: (1) both *PLT* and *CUC2* require downstream *YUC4* activity as ectopic overexpression of *PLT* proteins as well as of *CUC2* fails to trigger regeneration response in the *yuc4;yuc1* mutant; (2) reconstitution of *YUC4* expression under a heterologous promoter in the *plt* triple mutant as well as in the *cuc2-1D* mutant rescues the vascular regeneration defects; and (3) *PLT* proteins and *CUC2* act synergistically to activate *YUC4* transcription and repair the damaged tissues, which involves induction of vascular identity and proper polarisation of the polar auxin transporter PIN1.

Our study revealed a previously unrecognised role of *ANT* in vascular regeneration and demonstrated that a *PLT*-like gene from rice, a morphologically diverse grass species, could substitute the regeneration function of *Arabidopsis* *PLT* genes. These observations indicate that the activation of *PLT* gene promoters in response to mechanical injuries may be more important for the selection of regeneration-associated *PLT* genes than their protein sequence. In this context, it is relevant that distinct *PLT* transcription factors determine competence for regeneration in the root context (Durgaprasad et al., 2019).

In striking contrast to *in vitro* shoot regeneration cues (Kareem et al., 2015), *PLT3,5,7* do not act through the root stem cell regulators *PLT1*, *PLT2* and *WOX5* to initiate repair of damaged aerial tissues of a growing plant. Rather, *PLT* proteins act through *CUC2* by directly activating its expression (Fig. 6N). Interestingly, *PLT* genes and *CUC2* act in a feed-forward loop to activate the expression of the auxin biosynthesis gene *YUC4* (Fig. 6O). This circuit can act as a coherent feed-forward loop, which often serves as a signal persistence detector (Mangan et al., 2003), even though our analysis indicates that the regulatory logic at the promoter is not strictly an 'AND gate' (Alon, 2006). Regardless of the precise regulatory logic, the output of the circuit is the activation of *YUC4*. In that view, it is tantalising that the cellular defects associated with the malfunctioning of this circuit are the inability to redirect ground tissue cells to vascular identity and the inability to properly polarise PIN proteins. A regulatory feedback loop between auxin level, auxin flux and polarisation of auxin efflux carriers (PIN) has been proposed as a key regulatory mechanism of shoot branching, phyllotaxis and vascular tissue differentiation (Jönsson et al., 2006; Smith et al., 2006; Bayer et al., 2009; Schuetz et al., 2012; Mazur et al., 2016; Fujita and Kawaguchi, 2018). It is therefore conceivable that *PLT-CUC2*-dependent activation of *YUC4* activates this feedback loop to drive vascular regeneration in damaged growing leaves (Fig. 6O). In summary, our study reveals *PLT-CUC2* regulatory axis is specifically involved in controlling regeneration through induction of a local hormonal environment in response to injury.

## MATERIALS AND METHODS

### Plant materials

*Arabidopsis thaliana* ecotype Columbia (Col-0) was used as wild type in this study. The origins of the mutants used in the study, such as the double mutants *plt3;plt5-2*, *plt3;plt7* and *plt5-2;plt7* and the triple mutant *plt3;plt5-2;plt7* (Prasad et al., 2011), the double mutant *yuc4;yuc1* (Pinon et al., 2013), the single mutant *ant-4*, double mutant *ant-4;plt5-3* and the triple mutant *plt3;plt7;ant-4* (Křížek, 2015), the single mutants *cuc2-1D* (Larue et al., 2009) and *cuc2-3* (Hibara et al., 2006), and the double mutant *cuc1-5;cuc2-3* (Hibara et al., 2006), have been described previously. Translational fusion constructs of *PLT1::PLT1-vYFP*, *PLT2::PLT2-vYFP* (Mähönen et al., 2014), *PLT3::PLT3-vYFP*, *PLT5::PLT5-vYFP* and *PLT7::PLT7-vYFP* (Prasad et al., 2011) have been described previously. *35S::PLT5-GR*, *35S::PLT7-GR* (Prasad et al., 2011), *pCUC2::3XVENUS* and *35S::CUC2-3AT* (Kareem et al., 2015) have been described previously. Multisite gateway recombination cloning system

(Invitrogen) using the pCambia 1300 destination vector was used for cloning the translational fusion constructs, which were then introduced into the C58 *Agrobacterium* strain by electroporation and further transformed into wild-type or mutant *Arabidopsis* plants by the floral dip method (Clough and Bent, 1998) (see supplementary Materials and Methods for details on plasmid construction). *DR5rev::3XVENUS-N7* expression was examined in wild-type and *plt3;plt5-2;plt7* transgenic plants with the double marker *pDR5rev::3XVENUS-N7;PIN1::PIN1-GFP* line, which has been described previously (Pinon et al., 2013). In this study, only the YFP marker was analysed using a single YFP channel.

### Growth conditions

*Arabidopsis thaliana* seeds were surface sterilised with 70% ethanol and 20% bleach, followed by seven washes with sterile distilled water. Seeds were plated on half-strength Murashige-Skoog (MS) medium (pH 5.7) and grown vertically under 45  $\mu\text{mol}/\text{m}^2/\text{s}$  continuous white light at 22°C and 70% relative humidity.

### Regeneration assays

For wound-induced natural regeneration experiments, all plants and explants were grown on hormone-free half-strength MS agar medium (Sigma-Aldrich). To study wound repair and vascular regeneration in growing inflorescence stems, 3-week-old seedlings were selected. Using a sterile razor blade, the stem region between the rosette leaves and the first or second cauline leaves was subjected to either peeling of the tissue layers including epidermis and sub epidermal layers (inflorescence abrasion) or partial incision (inflorescence incision) through the vascular tissues under a dissection microscope (Zeiss Stemi 2000). The observations were recorded 4 days after wounding. For the leaf vascular regeneration assay, to maintain uniformity we injured a single leaf belonging to the first pair of rosette leaves of 5 dpv seedlings. Plants of same developmental stage were chosen for incision. Fine-pointed sterile tweezers (Dumont tweezer, Style 5) were used to make a sharp incision in the midvein at the basal part of the leaf blade. To avoid ambiguity, incisions made elsewhere were not scored. The incisions were made from the abaxial surface of the leaf to ensure precise injury to the midvein. The injured leaf was left connected to the growing parent plant and it was protected from any further damage. Vascular regeneration was analysed in the injured leaf 4 days post-incision. These leaves were cut at the petiole using Vannas straight scissors (Ted Pella, 1340) without causing additional damage to the leaf blade. The leaf tissue was cleared using chloral hydrate (Sigma-Aldrich) (see supplementary Materials and Methods for further details of decolourisation and tissue clearing) and brightfield images were obtained to assess the regeneration outcomes. When newly formed vascular strands (identified by the distinct morphology of end-to-end connected xylem elements) connected the cut ends of the midvein to form a D-shaped loop or connected the damaged midvein to a lateral vein, the outcomes were scored as successful regeneration (Fig. 2I,J). To study healing in response to wounding in excised organs (leaf/root), excised explants were collected from 9 dpv seedlings and placed on hormone-free MS agar medium. Upon excision, continuous dexamethasone (Sigma-Aldrich) induction was provided until the tenth day post-excision for samples collected from transgenic lines harbouring steroid-inducible constructs. The corresponding mock-treated samples were incubated on MS plates containing an equal proportion of DMSO (same volume as dexamethasone). The plates were kept in the dark for the first 24–32 h and later shifted to continuous light. All the plates of regeneration experiments were incubated vertically in a plant growth chamber maintained at 22°C and 70% relative humidity under 45  $\mu\text{mol}/\text{m}^2/\text{s}$  continuous white light.

### Microscopic imaging

Brightfield and confocal laser-scanning microscopy imaging were performed as described previously (Kareem et al., 2015). Brightfield images of vascular regeneration in incised leaves were acquired using the brightfield mode in a Leica TCS SP5 II inverted confocal microscope and an Olympus BX63F fluorescence microscope after clearing the leaf sample (see supplementary Materials and Methods for details of decolourisation and tissue clearing). Confocal imaging of leaves and thick samples were performed using a Leica TCS SP5 II upright microscope and a Zeiss LSM

880 confocal laser-scanning microscope. Brightfield images acquired using Leica M205 FA fluorescence stereo microscope and confocal microscopes were adjusted for brightness and contrast. For confocal imaging, the cell boundaries of inflorescence stem samples were stained using 10  $\mu\text{g}/\text{ml}$  propidium iodide (Sigma-Aldrich). Images were acquired using 10 $\times$  air, 20 $\times$  oil immersion, 20 $\times$  air and 40 $\times$  oil immersion objectives. The projection view of the images was reconstructed from the z-stacks with Leica LAS-AF software and Zeiss ZEN blue softwares. Images were compiled using Adobe Photoshop CS6 and Adobe Photoshop CC 2015. All image panels represent z-stacks unless mentioned. The area of callus formation at the cut end of detached organs was measured using ImageJ software.

### qRT-PCR

Total RNA was extracted from samples (see supplementary Materials and Methods for further details of sample preparation) using the Nucleospin Plant RNA extraction kit (MN) and subjected to on-column DNase treatment according to the manufacturer's guidelines. cDNA was synthesised from 1  $\mu\text{g}$  total RNA using a High-Capacity cDNA Reverse Transcription kit (Applied Biosystems). qRT-PCR was performed in a 25  $\mu\text{l}$  reaction volume containing 12.5  $\mu\text{l}$  SYBR Green PCR master mix (Takara Bio), 100 nM gene-specific primers (Table S1) and 100 ng cDNA in a CFX96 Touch Real-Time PCR Detection System. All reactions were performed with RNA derived from three independent biological replicates. Each biological sample was tested in technical triplicate. Data were normalised to *ACTIN2* (*ACT2*). The transcript level in the control was normalised to 1. The expression of the gene of interest is represented with respect to the control (as performed by Kareem et al., 2015). The relative gene expression is represented as fold-change value by calculating  $-\Delta\Delta C_T$ .

### Luciferase assay

The luciferase assay was performed as described by Díaz-Triviño et al. (2017). Healthy *Nicotiana benthamiana* plants (3–4 weeks old) grown under long-day conditions (16 h light, 8 h dark) were used for agroinfiltration. The primers used for cloning are listed in Table S2. Competent cells of the C58 strain of *Agrobacterium tumefaciens* were used for the infiltration.

### ChIP-qPCR analysis

ChIP was performed by following the protocol as described by Yamaguchi et al. (2014) (see supplementary Materials and Methods for a brief description). ChIP-qPCR was performed using SYBR Premix (Clontech) to determine PLT5 protein occupancy on the *CUC2* promoter region. The relative fold enrichment of *CUC2* DNA was calculated by computing the enrichment in *PLT5::PLT5-vYFP* relative to *plt3;plt5-2;plt7*. *ACTIN7* (*ACT7*) was used to normalise the results between the samples. The ChIP-qPCR reactions were performed in triplicate. The primers used for ChIP-qPCR are listed in Table S3.

### Statistical analysis

Pearson's  $\chi^2$  test (regeneration assay analysis), Welch's two-sample *t*-test (qRT-PCR data analysis), Mann–Whitney *U* one-tailed test (luciferase assay) and Kruskal–Wallis  $\chi^2$  test (comparing the number of closed vein loops) were used for data analysis. The Holm–Bonferroni correction was performed for multiple analysis when using Pearson's  $\chi^2$  test. R programming was used for the statistical analyses.

### Acknowledgements

We thank Dr Philip N. Benfey, Dr Elliot Meyerowitz and Dr Ottoline Leyser for their valuable suggestions on the early draft of the manuscript. We are grateful to Dr Charles Melnyk and Dr Utpal Nath for all the discussions and help. *pG10-90::vYFP* control vector for luciferase assay was received from Menno Pijnenburg. We are thankful to M. R. Krishna Prashanth for assistance with schematics and figures, B. Subhiksha for cloning *AtPLT5::OsPLT2-vYFP* and Sajesh Vijayan for help with the statistical analyses.

### Competing interests

The authors declare no competing or financial interests.

### Author contributions

Conceptualization: D.R., A.K., K.P.; Methodology: D.R., A.P.S., A.K., M.A., V.V., A.T., M.K., D.V., A.N.L., V.W., K.P.; Formal analysis: D.R., A.P.S., A.K., M.A., A.T.,

A.S., M.K.M., V.W., B.S., K.P.; Investigation: A.K., K.P.; Resources: A.K., M.G.S., E.S., B.A.K., I.E., A.P.M., B.S., K.P.; Writing - original draft: D.R., A.K.; Writing - review & editing: D.R., A.P.S., A.T., B.S., K.P.; Visualization: D.R., A.P.S., A.K., K.P.; Supervision: K.P.; Funding acquisition: K.P.

## Funding

K.P. acknowledges grants from the Department of Biotechnology (DBT), Government of India [grant BT/PR12394/AGIII/103/891/2014] and Department of Science and Technology, Science and Engineering Research Board (DST-SERB), Government of India [grant EMR/2017/002503/PS] and also acknowledges the Indian Institute of Science Education and Research Thiruvananthapuram (IISER-TVM) for infrastructure and financial support. D.R. acknowledges a University Grants Commission (UGC) fellowship. A.P.S., V.V. and A.T. are recipients of Council of Scientific and Industrial Research (CSIR) fellowships. A.K. was supported by an Indian Institute of Science Education and Research Thiruvananthapuram fellowship. M.A. acknowledges Department of Biotechnology (DBT), Ministry of Science and Technology, Government of India for granting the DBT-Post Doctoral Fellowship (DBT-RA Program). D.V. was a recipient of an Innovation in Science Pursuit for Inspired Research scholarship, and A.N.L. was supported by a Kishore Vaigyanik Protsahan Yojana scholarship of the Department of Science and Technology, Government of India. A.P.M. acknowledges grants from the Academy of Finland [grants 266431 and 271832]. I.E. acknowledges the Israel Science Foundation [ISF966/17] and the Howard Hughes Medical Institute [International Research Scholar Grant 55008730]. B.A.K. was supported by a grant from National Science Foundation (NSF) [grant NSFIO5135442]. E.S. was supported by Discovery Grants of the Natural Science and Engineering Research Council of Canada [grants NSERC RGPIN-2016-04736 and NSERC RGPAS 492872-2016].

## Data availability

All raw data associated with this article have been deposited in Mendeley (<https://data.mendeley.com/datasets/mwyxw4v63h/draft?a=e64505aa-564b-4127-9d0c-afc900810544>).

## Supplementary information

Supplementary information available online at <http://dev.biologists.org/lookup/doi/10.1242/dev.185710.supplemental>

## Peer review history

The peer review history is available online at <https://dev.biologists.org/lookup/doi/10.1242/dev.185710.reviewer-comments.pdf>

## References

- Alon, U. (2006). *An Introduction to Systems Biology*. New York: Chapman and Hall/CRC.
- Asahina, M., Azuma, K., Pitaksaringkarn, W., Yamazaki, T., Mitsuda, N., Ohme-Takagi, M., Yamaguchi, S., Kamiya, Y., Okada, K., Nishimura, T. et al. (2011). Spatially selective hormonal control of RAP2.6L and ANAC071 transcription factors involved in tissue reunion in Arabidopsis. *Proc. Natl. Acad. Sci. USA* **108**, 16128-16132. doi:10.1073/pnas.1110443108
- Bayer, E. M., Smith, R. S., Mandel, T., Nakayama, N., Sauer, M., Prusinkiewicz, P. and Kuhlemeier, C. (2009). Integration of transport-based models for phyllotaxis and midvein formation. *Genes Dev.* **23**, 373-384. doi:10.1101/gad.497009
- Bisborough, G. D., Runions, A., Barkoulas, M., Jenkins, H. W., Hasson, A., Galinha, C., Laufs, P., Hay, A., Prusinkiewicz, P. and Tsiantis, M. (2011). Model for the regulation of Arabidopsis thaliana leaf margin development. *Proc. Natl. Acad. Sci. USA* **108**, 3424-3429. doi:10.1073/pnas.1015162108
- Clough, S. J. and Bent, A. F. (1998). Floral dip: a simplified method for Agrobacterium-mediated transformation of Arabidopsis thaliana. *Plant J.* **16**, 735-743. doi:10.1046/j.1365-3113x.1998.00343.x
- Díaz-Triviño, S., Long, Y., Scheres, B. and Blilou, I. (2017). Analysis of a plant transcriptional regulatory network using transient expression systems. In *Methods in Molecular Biology* (ed. K. Kaufmann and B. Mueller-Roeber), pp. 83-103: Humana Press.
- Donner, T. J., Sherr, I. and Scarpella, E. (2009). Regulation of preprocambial cell state acquisition by auxin signaling in Arabidopsis leaves. *Development* **136**, 3235-3246. doi:10.1242/dev.037028
- Du, Y. and Scheres, B. (2017). PLETHORA transcription factors orchestrate de novo organ patterning during Arabidopsis lateral root outgrowth. *Proc. Natl. Acad. Sci. USA* **114**, 11709-11714. doi:10.1073/pnas.1714410114
- Durgaprasad, K., Roy, M. V., Venugopal, M. A., Kareem, A., Raj, K., Willemsen, V., Mähönen, A. P., Scheres, B. and Prasad, K. (2019). Gradient expression of transcription factor imposes a boundary on organ regeneration potential in plants. *Cell Rep.* **29**, 453-463.e3. doi:10.1016/j.celrep.2019.08.099
- Efroni, I., Mello, A., Nawy, T., Ip, P.-L., Rahni, R., DelRose, N., Powers, A., Satija, R. and Birnbaum, K. D. (2016). Root regeneration triggers an embryo-like sequence guided by hormonal interactions. *Cell* **165**, 1721-1733. doi:10.1016/j.cell.2016.04.046
- Flaishman, M. A., Loginovsky, K. and Lev-Yadun, S. (2003). Regenerative xylem in inflorescence stems of Arabidopsis thaliana. *J. Plant Growth Regul.* **22**, 253-258. doi:10.1007/s00344-003-0030-y
- Fujita, H. and Kawaguchi, M. (2018). Spatial regularity control of phyllotaxis pattern generated by the mutual interaction between auxin and PIN1. *PLoS Comput. Biol.* **14**, e1006065. doi:10.1371/journal.pcbi.1006065
- Galliot, B., Crescenzi, M., Jacinto, A. and Tajbakhsh, S. (2017). Trends in tissue repair and regeneration. *Development* **144**, 357-364. doi:10.1242/dev.144279
- Hibara, K.-I., Karim, M. R., Takada, S., Taoka, K.-I., Furutani, M., Aida, M. and Tasaka, M. (2006). Arabidopsis CUP-SHAPED COTYLEDON3 regulates postembryonic shoot meristem and organ boundary formation. *Plant Cell* **18**, 2946-2957. doi:10.1105/tpc.106.045716
- Ikeuchi, M., Ogawa, Y., Iwase, A. and Sugimoto, K. (2016). Plant regeneration: cellular origins and molecular mechanisms. *Development* **143**, 1442-1451. doi:10.1242/dev.134668
- Ikeuchi, M., Shibata, M., Rymen, B., Iwase, A., Bågman, A.-M., Watt, L., Coleman, D. S., Takahashi, T., Ahnert, S. E. et al. (2018). A gene regulatory network for cellular reprogramming in plant regeneration. *Plant Cell Physiol.* **59**, 765-777. doi:10.1093/pcp/pcy013
- Jönsson, H., Heisler, M. G., Shapiro, B. E., Meyerowitz, E. M. and Mjolsness, E. (2006). An auxin-driven polarized transport model for phyllotaxis. *Proc. Natl. Acad. Sci. USA* **103**, 1633-1638. doi:10.1073/pnas.0509839103
- Kareem, A., Durgaprasad, K., Sugimoto, K., Du, Y., Pullanmackal, A. J., Trivedi, Z. B., Abhayadev, P. V., Pinon, V., Meyerowitz, E. M., Scheres, B. et al. (2015). PLETHORA genes control regeneration by a two-step mechanism. *Curr. Biol.* **25**, 1017-1030. doi:10.1016/j.cub.2015.02.022
- Kokla, A. and Melnyk, C. W. (2018). Developing a thief: haustoria formation in parasitic plants. *Dev. Biol.* **442**, 53-59. doi:10.1016/j.ydbio.2018.06.013
- Krizek, B. A. (2015). AINTEGUMENTA-LIKE genes have partly overlapping functions with AINTEGUMENTA but make distinct contributions to Arabidopsis thaliana flower development. *J. Exp. Bot.* **66**, 4537-4549. doi:10.1093/jxb/erv224
- Larue, C. T., Wen, J. and Walker, J. C. (2009). A microRNA-transcription factor module regulates lateral organ size and patterning in Arabidopsis. *Plant J.* **58**, 450-463. doi:10.1111/j.1365-3113X.2009.03796.x
- Mähönen, A. P., Ten Tusscher, K., Siligato, R., Smetana, O., Díaz-Triviño, S., Salojärvi, J., Wachsmann, G., Prasad, K., Heidstra, R. and Scheres, B. (2014). PLETHORA gradient formation mechanism separates auxin responses. *Nature* **515**, 125. doi:10.1038/nature13663
- Mangan, S., Zaslaver, A. and Alon, U. (2003). The coherent feedforward loop serves as a sign-sensitive delay element in transcription networks. *J. Mol. Biol.* **334**, 197-204. doi:10.1016/j.jmb.2003.09.049
- Marhava, P., Hoermayer, L., Yoshida, S., Marhavý, P., Benková, E. and Friml, J. (2019). Re-activation of stem cell pathways for pattern restoration in plant wound healing. *Cell* **177**, 957-969.e13. doi:10.1016/j.cell.2019.04.015
- Mazur, E., Benková, E. and Friml, J. (2016). Vascular cambium regeneration and vessel formation in wounded inflorescence stems of Arabidopsis. *Sci. Rep.* **6**, 33754. doi:10.1038/srep33754
- Melnyk, C. W., Schuster, C., Leyser, O. and Meyerowitz, E. M. (2015). A developmental framework for graft formation and vascular reconnection in Arabidopsis thaliana. *Curr. Biol.* **25**, 1306-1318. doi:10.1016/j.cub.2015.03.032
- Nikovics, K., Blein, T., Peaucelle, A., Ishida, T., Morin, H., Aida, M. and Laufs, P. (2016). The balance between the MIR164A and CUC2 genes controls leaf margin serration in Arabidopsis. *Plant Cell* **18**, 2929-2945. doi:10.1105/tpc.106.045617
- Ohashi-Ito, K., Oguchi, M., Kojima, M., Sakakibara, H. and Fukuda, H. (2013). Auxin-associated initiation of vascular cell differentiation by LONESOME HIGHWAY. *Development* **140**, 765-769. doi:10.1242/dev.087924
- O'Malley, R. C., Huang, S.-C., Song, L., Lewsey, M. G., Bartlett, A., Nery, J. R., Galli, M., Gallavotti, A. and Ecker, J. R. (2016). Cistrome and epistrome features shape the regulatory DNA landscape. *Cell* **165**, 1280-1292. doi:10.1016/j.cell.2016.04.038
- Pinon, V., Prasad, K., Grigg, S. P., Sanchez-Perez, G. F. and Scheres, B. (2013). Local auxin biosynthesis regulation by PLETHORA transcription factors controls phyllotaxis in Arabidopsis. *Proc. Natl. Acad. Sci. USA* **110**, 1107-1112. doi:10.1073/pnas.1213497110
- Pitaksaringkarn, W., Matsuoka, K., Asahina, M., Miura, K., Sage-Ono, K., Ono, M., Yokoyama, R., Nishitani, K., Ishii, T., Iwai, H. et al. (2014). XTH20 and XTH19 regulated by ANAC071 under auxin flow are involved in cell proliferation in incised Arabidopsis inflorescence stems. *Plant J.* **80**, 604-614. doi:10.1111/tpj.12654
- Prasad, K., Grigg, S. P., Barkoulas, M., Yadav, R. K., Sanchez-Perez, G. F., Pinon, V., Blilou, I., Hofhuis, H., Dhonukshe, P., Galinha, C. et al. (2011). Arabidopsis PLETHORA transcription factors control phyllotaxis. *Curr. Biol.* **21**, 1123-1128. doi:10.1016/j.cub.2011.05.009
- Radhakrishnan, D., Kareem, A., Durgaprasad, K., Sreeraj, E., Sugimoto, K. and Prasad, K. (2018). Shoot regeneration: a journey from acquisition of competence to completion. *Curr. Opin. Plant Biol.* **41**, 23-31. doi:10.1016/j.pbi.2017.08.001
- Reinhardt, D., Frenz, M., Mandel, T. and Kuhlemeier, C. (2003). Microsurgical and laser ablation analysis of interactions between the zones and layers of the tomato shoot apical meristem. *Development* **130**, 4073-4083. doi:10.1242/dev.00596



- Sachs, T.** (1968). On the determination of the pattern of vascular tissue in peas. *Ann. Bot.* **32**, 781-790. doi:10.1093/oxfordjournals.aob.a084249
- Sachs, T.** (1969). Polarity and the induction of organized vascular tissues. *Ann. Bot.* **33**, 263-275. doi:10.1093/oxfordjournals.aob.a084281
- Sachs, T.** (1981). The control of the patterned differentiation of vascular tissues. In *Advances in Botanical Research* (ed. H. W. Woolhouse), pp. 151-262: Academic Press.
- Sachs, T.** (1991). *Pattern Formation in Plant Tissues*. New York: Cambridge University Press.
- Scarpella, E., Francis, P. and Berleth, T.** (2004). Stage-specific markers define early steps of procambium development in Arabidopsis leaves and correlate termination of vein formation with mesophyll differentiation. *Development* **131**, 3445-3455. doi:10.1242/dev.01182
- Scarpella, E., Marcos, D., Friml, J. and Berleth, T.** (2006). Control of leaf vascular patterning by polar auxin transport. *Genes Dev.* **20**, 1015-1027. doi:10.1101/gad.1402406
- Schuetz, M., Smith, R. and Ellis, B.** (2012). Xylem tissue specification, patterning, and differentiation mechanisms. *J. Exp. Bot.* **64**, 11-31. doi:10.1093/jxb/ers287
- Smetana, O., Mäkilä, R., Lyu, M., Amiryousefi, A., Sánchez Rodríguez, F., Wu, M. F., Solé-Gil, A., Leal Gavarrón, M., Siligato, R., Miyashima, S. et al.** (2019). High levels of auxin signalling define the stem-cell organizer of the vascular cambium. *Nature* **565**, 485-489. doi:10.1038/s41586-018-0837-0
- Smith, R. S., Guyomarc'h, S., Mandel, T., Reinhardt, D., Kuhlemeier, C. and Prusinkiewicz, P.** (2006). A plausible model of phyllotaxis. *Proc. Natl. Acad. Sci. USA* **103**, 1301-1306. doi:10.1073/pnas.0510457103
- van den Berg, C., Willemsen, V., Hage, W., Weisbeek, P. and Scheres, B.** (1995). Cell fate in the Arabidopsis root meristem determined by directional signalling. *Nature* **378**, 62-65. doi:10.1038/378062a0
- Wenzel, C. L., Schuetz, M., Yu, Q. and Mattsson, J.** (2007). Dynamics of MONOPTEROS and PIN-FORMED1 expression during leaf vein pattern formation in Arabidopsis thaliana. *Plant J.* **49**, 387-398. doi:10.1111/j.1365-313X.2006.02977.x
- Xu, J., Hofhuis, H., Heidstra, R., Sauer, M., Friml, J. and Scheres, B.** (2006). A molecular framework for plant regeneration. *Science* **311**, 385-388. doi:10.1126/science.1121790
- Yamaguchi, N., Winter, C. M., Wu, M. F., Kwon, C. S., William, D. A. and Wagner, D.** (2014). PROTOCOL: Chromatin Immunoprecipitation from Arabidopsis Tissues. *Arabidopsis Book* **12**, e0170. doi:10.1199/tab.0170
- Zhou, W., Lozano-Torres, J. L., Blilou, I., Zhang, X., Zhai, Q., Smant, G., Li, C. and Scheres, B.** (2019). A jasmonate signaling network activates root stem cells and promotes regeneration. *Cell* **177**, 942-956.e14. doi:10.1016/j.cell.2019.03.006